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FINAL REPORT

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BETHANY COLLEGE BETHANY, V. VA.

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A STUDY OF THE PREPARATION AND ELECTRICAL PROPERTIES OF CERTAIN CLAY FILM DIFLECTRICS

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STATE FOR OF THE PROBLEM

An experimental study was made of bentonite clay films
(alsifilms) with respect to their physical, (4, 5, 6, 7, 8),
electrical and colloidal properties. The variables considered
were: bentonites from various sources, bentonites of various
capacities for base exchange, bentonites of various particle size
distributions, various base exchange agents, and subsequent heat
cures or pressure and heat cures on the base exchanged films.

CHOICE OF CLAY

The choice of the clay to be used in this study proved to be very important because of the differences in the range of particle sizes of samples from various sources. To make a successful alsifilm there had to be no clay particles greater in size than five microns and most had to be less than that size. Some clays occur naturally with larger proportions of these small particles than do others. Bentonites are favorable in this respect in contrast to halloysite, for example, which has too small a percentage of the particles in the desired size range. Not all bentonites have the same range of particle sizes, however, as is shown in Table I.

Another basis of comparison and contrast of these various clays, in addition to particle size, is the chemical composition. Not much correlation has been drawn between chemical composition and film properties other than the color of the film. All films prepared in this study had a light tan color before base exchange and heat cure. This color was attributed to the presence of iron oxides in the bentonite. This belief was later substantiated

TABLE I

Type of Clay	thon 74	Percer Larger than 44 micron	Less then 44	of Vario	Less then	3-5	0.5-5	0.5-l micron	Less than 0.5 micron
Volcley (1)		0.1-4		2-3	94			4-5	80- 90
Average Colloidel Bentonite (1)		3-4		5-10				20- 30	60- 70
Non-bento plastic clay (1)	nîte	15- 20		30- 50				25- 40	15- 35
Fisher (B-237/2) (3)			95						
Bennett Clay (15)	3-4								
Panther Creek (1)					7 0		3 5		35
Wyodak micro- mesh (17)						99			

Attacley (2) Average particle size by air permeation method is 1-2 microns

Please Note: 1 micron is approximately 1/25000 inch; 50 microns dismeter--smallest single particle visible to the human eye

in part by the light colored clay obtained from the base exchange experiments when iron was replaced by hydrogen. From the point of view of low iron content, Bennett clay (Table II) seemed ideal. However, the average particle size of this clay was entirely too large to permit the preparation of suspensions which could be centrifuged. Most of the clay settled out in the 24 hour period before the actual centrifuging was begun. Table II gives the approximate composition of a number of the bentonite clays.

The final selection of clay used in this work was Wyodak micro-mesh and American Colloid Volclay. In oreliminary work, a bentonite (B-237/2) furnished by Fisher Scientific Co.. my was used.

CENTRIFUCING

A fractionation of the clay in water suspension was necessary in order to concentrate the particles in the desired size range from the colloidal clays selected. This operation was carried out with a Sharoles Super Centrifuge (Type M-41-24). Various slip feeds and RPM's of the bowl were tried before the best procedure was found.

In the first separations of Fisher Scientific bentonite, the solid material left in the centrifuge bowl after the second pass was suspended again in distilled water, boiled to the gel stage, and laid down in the film forming step. In this procedure the slip feed of the first bass was 90 ml. per minute at a bowl speed of 12,000 RPM. For the second pass the slip feed was 20 ml, per minute and the bowl speed was 16,000 RPM. Although a large percentage of the perticles was in the size range less than 0.07 micron in diameter. enough of the particles were greater than that size to give a very course and brittle-film.

TABLE II
CHEMICAL ANALYSIS OF VARIOUS CLAYS

Chemical Analysis	Volclay	Creek	Artosite (15)	Cley	micro-	Fisher (3)	Attapulgus Clay (2)	
Percent		Bentonite (1)		(15)	mesh (17)		• ,	
Sio _s	64.32	64.17	60.16	71.00	65.17	63.20	68.43	
Al ₂ O ₃	20.74	17.14	19.88	13.00	16.64	20.54	12.56	
Fe ₂ 0 ₅	3.03	4.81	2.97	0.85	4.91	3.25	4.94	
FaO	0.46		0.54			0.26		
CaO	0.52	1.48	0.67	1.20	1.39	1.30	1.67	
MgO	2.30	3.90	2.20	2.04	3.78	2.49	9.82	
NegO	2.59	0.21	2.60	2.30	0.19	2.16	0.25	
K ₂ 0	0.39	0.48	0.40	0.60	0.46	0.50	0.70	
Essentia Moisture			4.91			5.06		
Ignition Loss		7.78	5 50	0.15		2 22		
		7.70	5.58	9.13	7.44	7.19	12.10	
pH (H ₂ 0 suspen- sion)	8.5~ 9.5	6-6.5	9.2	5.1			7-7.6	

Further refinements in the centrifuging procedure were made.
Four or six passes were run on an original two percent suspension of the bentonite in distilled water. The suspension stood for 24 hours after its preparation before the centrifuging began. The colloidal gel was made from the liquid pass of the last separation and not from the larger particles in the bowl. Both the Wyodak and the American Colloid clays were treated in this way.

Hauser and Reed (14) have worked out a graphical method for correlating the PPH and the slip feed of a Sharples Super Centrifuge with the particle sizes settled out under these conditions. These particle sizes along with the bowl speeds and slip feeds employed in the four and the six step procedures are given in Tables III and IV.

TABLE III

FOUR SMEP CTUM ITUCING PROCEDURE WAS AS FOLLOWS:

Bowl Speed (RPM)	Slip Feed (ml. per minute)	Particle size in the pass
6000	100	Less than 0.4
10000, later 8000	70, later 90	0.2 micron, later 0.3 micron
10000	70	0.2 micron and less
10000	70	0.2 micron and less

Substantial evaporation of the last pass in the four step procedure listed in Table III gave a gel which resulted in a very thin film (0.001 inch in thickness) of fair strength. The smaller colloidal particles in the last pass of the six step procedure gave even a tougher film. The degree of texture smoothness and strength of the film was in indirect ratio to the size of the colloidal particles employed.

TABLE IV
SIX STEP CONTRIBUGING PLOCEDURE WAS AS FOLLOWS:

Bowl Speed (RPM)	Slip Feed (ml. per minute)	Particle size in the pass
6000	100	Less than 0.4 micron
8000	90	0.3 micron and less
10000	70	0.2 micron and less
15000	40	0.1 micron and less
23000	40	Less than 0.07 micron
23000	40	Less than 0.07 micron

This method of using the liquid pass from the last run instead of the larger solid particles from the bowl in preparing the gel was in direct contrast to the methods of Hauser (5) and was similar to the method used by Aircraft-Varine Products Inc. (16)

EVAPORATION AND GEL FORMATION

Portions of the last bass from the centrifuting process were put into two liter beakers and evaporated on electric hot plates with constant mechanical stirring. Concentration continued in each case until the system had a viscosity very similar to that of a bentonite dispersion in water which, when tested by a Stormer Viscosimeter with a 99.26 gram weight attached, required 22.75 seconds for 100 revolutions at 30 degrees Centrigrade. A suspension of this viscosity gave a good gel.

Although the phenomenon of gelation is fairly common, it is still not thoroughly understood. Hauser and LeBeau (9) were able to prepare gels at concentrations of disperse phase as low as 0.05 percent. No slighment or grouping of the disperse phase was observed. It does not appear that an interlocking of long chain molecules need occur in order for gelation to take place.

FILM FORMATION

Various techniques have been employed in this operation.

D. W. Light (10) added a resin to a colloidel clay suspension. The mixed suspension was evaporated from an amalgam surface. The plasticizer was burned cut with oxygen.

Mershall (11, 12) formed films from electrodialyzed suspensions by evaporating them from copper or celluloid surfaces.

In this study many surfaces were tried for the evaporation step.

Among others the gelatinous clay was spread on tin, copper, paraffin covered glass and aluminum surfaces. The films stuck to the tin, copper and aluminum surfaces and the paraffin coating on glass melted at the temperature necessary for an advanced rate of drying.

A very satisfactory wax costed glass surface was developed at the suggestion of Mr. F. A. Wells. (16) Johnson's "Paste Wax" was first dissolved in benzene (10 to 15 percent solution by weight) and the solution was sprayed on the glass surface in three or four successive applications. The solvent evaporated leaving only the thin wax coating which did not melt at temperatures of 50 degrees Centigrade. A deep scoring or scratch was placed as border around the glass surface area. This scoring served to hold the film in place at the edges later during the drying operation and to prevent premature peeling of the film from the surface.

The freshly boiled gel suspension, while still warm (50 to 60 degrees Centigrade), was boured on the wax costed surfaces (four by six inches) and was spread by a stainless steel scraping bar to exactly 0.075 inch thickness. It was bointed out by Mr. Wells (16) that initial wet layers much in excess of 0.075 inch thickness will not dry to tough self supporting films. They generally crack into many pieces, are not

solf-supporting and are very brittle.

The next stop, the drying of the film, was a critical one. Air drying at room temperature was first tried, but that procedure, requiring about a wook, was too time consuming. Next, infra-rod heating either from the top or the bottom was tried, but in every instance extreme cracking of the film occurred near the end of the operation and the film would stick to the class surface. Finally, a forced air drying cabinet was built and drying in a stream of dust free air at 50 degrees Contigrade for about four hours gave the desired results. If the film was left in the cabinet for longer periods, however, it would weld to the glass surface and be impossible to remove in one piece. In the drying procedure the gel layer shrank to 1/75 th of its original wet thickness, but no cracking or surface separation was noted. Thicker films could best be prepared by laminating a number of these thin dry films under treatment of heat and pressure.

Some work was also done on self-supporting clay films reinforced with asbestos fibers and cellulose shreds. One to two percent by weight of shredded asbestos or filter paper was stirred vigorously into the gelation mixture of the clay. The rest of the procedure of film forming and drying was carried out as above. These films were tough but were not uniform in texture. This approach was abandoned when strong self-supporting films containing only bentonite were propared.

BASE EXCHANGE

A clay is really a very complex anion associated with soveral cations. In acueous suspension these cations dissociate and the clay micelle may be thought of as a dissociated electrolyte. One cation of the clay may then be exchanged for another just as a sodium soap will exchange its cation for calcium in hard water.

Not all clays will beso exchange to the same extent. Also, after some cations are replaced by others, such as calcium for sodium, the clay greatly loses its swelling properties in water. It has been established that the order of preference of the clay anion for various cations is calcium, magnesium, potassium, hydrogen, sodium, lithium.(1)

It was hoped that the larger cations, approximating in size the dimonsions between atems in the montmorillenite minerals, would enter these spaces in the base exchange and be locked into them during the curing process. From this viewpoint larger cations would be preferred to smaller ones in the base exchange step. Large organic radicals were also of interest in base exchange work, but they would not withstand the heat cure process which appears to be necessary for reducing the power factor in these dielectric films to a desirable value.

Table V.gives data on the exchangaable metallic bases in milliequivalents per 100 grams of some clays. (1)

		TABLE			
Cations	Volclay	tonites Panther Creek			Kentucky
*******	MAOBIUS	Mississippi	<u>abeven</u>	Fire Clay	Ball Clay
calcium	22.0	64.7	51.5	5.2	7.1
sodium	85.5	0.4	33.6	0.4	0.5
potassium	5.0	2.8	1.1	0.5	0.4
me gnesium	1.0	1.0	19.1	1.3	3.8

various base exchange techniques for bentonite were investigated using saturated barium chloride solutions with the clay. By one method the dried self-supporting transparent film was suspended in the saturated solution of the salt for a period of two or three hours. On removal of the film from solution, any cryatels of the salt adhering to the film were washed off and the film was dried at 50 degrees Centigrade.

This procedure was tenerally unsatisfactory. In the base exchange process by this method the self-supporting film usually had a tendency to disintegrate. If the film was not destroyed in this manner in the base exchange step, then in the drying process in the forced sir cabinet at 50 degrees Centigrade the film would curl and crack.

Successful bace exchange reactions were carifed out on the concentrated clay suspension from the sixth centrifuge pass in a Bredfield electrodialysis apparatus. Over a period of four hours the hydrogen clay formed in the central cell and the basic metal ions migrated to the cathode compartment. The voltage was 115 volta D. C., the amberage was 0.06, the rate of flow of distilled water through the anode compertment was 48 ml. per minute and through the cathode compartment was 30 ml. per minute. The scid clay was concentrated to s gel and a film was laid down. In order to introduce the desired cetion into the clay, this film was troated with a seturated barium hydroxide solution while still atteched to the wax costed glass plate. Another method was to add the barium hydroxide solution directly to the acid clay suspension, evaporate to the gel stage and then form the film. Either method worked satisfactorily. The only disadvantage was the small amount of scid clay that could be formed at any one time in the electrodialysis opporatus.

Another attempt was made to base exchange the cations before the film was poured. In this procedure dry Volclay was placed in a vertical glass column one inch in diameter and concentrated hydrochloric acid was poured through the packed column of clay. The acid flowed through extremely slowly under gravity. A definite displacement of ferric, calcium and magnesium ions from the clay by the hydrogen ion of the

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acid was indicated by qualitative test run on the filtrate. The resulting hydrogen clay was washed and dried and was then used in making a two percent acusous suspension in distilled water. This material was centrifused six times by the regular procedure. The sixth pass did not yield a satisfactory gel when it was concentrated.

For some time base exchange of the hydrogen clay with the appropriate metal hydroride seemed to be the ideal method of attack. It eliminated the need for removal of any foreign cations or anions since the hydrogen of the clay and the hydroxyl of the base gave only water which was lost in the drying step.

As suggested in work done by Hauser (6) berium chloride in various concentrations was thoroughly mixed with the concentrated gel. Films were then formed in the regular way. The mixture containing one percent berium chloride gave the best film of this type. The film was opaque and brittle. Films containing over one percent berium chloride added in this way were extremely brittle and contained crystals of berium chloride in the film surface. It does not appear that this process can be called base exchange, however, since no provision has been made in the procedure either for the removal of the cations (sodium, potassium, magnesium, calcium) naturally occurring in the clay or the anion (chloride, in this case) from the base exchange agent.

The most satisfactory way found in these studies to carry out the base exchange on a fairly large scale was to operate on the raw film after it had been dried in the cabinet and before it had been stripped from the glass plate on which it was formed. The plate with the film attached was immersed in a saturated solution of the barium

exchange, the film, still attached to the plate, was rinsed with distilled water to remove any surface adsorbed crystals or foreign ions, dried and then stripped from the plate and cut into pieces for testing. The disadvantage of this method, of course, was that the exchange could take place from only one side of the film. The advantage was, however, that the film remained intact and flat in the saturated salt solution, a situation that did not exist when the detached self-supporting film was introduced directly into an acueous solution.

Saturated solutions of the base exchange agents must be employed in order to prevent the partization of the film when introduced into the acucous solution.

Certain information concerning the base exchange agents used is given in Table VI.

CURING PROCESS

This trestment involved host or hest and pressure applied to the film after the base exchange step. It was hoped that the cure would be an effective method of locking in the larger exchange cations between the atoms in the bentonite grouping of atoms. The process appeared necessary to lower the power factor to a desirable value in the completed film.

Curing treatments for varying lengths of time were employed using heat treatments up to 550 degrees Centigrade and heat and pressure treatments with a Carver electrically heated press up to 175 degrees Centigrade and 8.5 tons. These specific conditions are listed in Table VII.

The standard procedure which was finally adopted was a gradual heating from room temperature to 550 degrees Centigrade over a period

TABLE VI

COMPOUNDS USED IN SATURATED AQUEOUS SOLUTIONS IN THE BASE

THICHANGE STEP

Compound	Sou	rce	Crade	Code	Formula
cerium nitrate	Beker an	d Adamson	Puri- fied	1560	
cobalt chlorids	J. m. Ba	ker	C. P.		CoCla.6HgO
cadmium acetate	Baker and	d Adamson	Reagent	1481	Cd(CH3C00)a.2Ha0
e lum	•• ••	• •	••	1238	Alm (SO4)3.KmSO4 .24HmO
thorium nitrate	•• ••	• •	••	2386	Th(NO ₃) _{4.4} H ₂ O
uranium acetate	••	••	••	2404	UO2 (CH2COO)2.2H2O
chronium sulfate	Eimer en	d Amend	C. P.	C-338	Cr ₂ (SO ₄) ₃ .aq.
sodium stennate	Baker an	d Adamson	Reagent	2292	Na2SnO3.3H2O
berium hydroxide	Merck		••	37211	Ba (OH) 2. BH O
lesd acetate	Eimer en	d Amend	C. P.		Pb(CH ₅ C00) ₂ .3H ₂ 0
berium chloride	J. T. Ba	ker	C: P.		BaClg.2HgO
calcium chloride	Eimer an	d Amend	C. P.	C-78	CaCls.6Hs0
calcium hydroxid	е	• •	Realent		Ca(OH) _s
mercurous nitrat	e Baker a	nd Adamso	n	1982	Egg (NO3)2.2H20
mercuric nitrate	<i>i</i>	• •	• •	1968	Hg(NO ₃)2.H2O

CLAY FILM DIELECTRICS

TABLE VII

Sample Number	Clay Used	Remark Preper	s on Film ation	Base Exchi Agent	ange
1	American Colloid Volclay	poured thickn dried	uspension to 2 mm. ess, then at 50 deg. cabinet	Saturated solution	BeClg
2	• •	• •	• •	• •	• •
3	• •	• •	• •	none	
4	• •	• •	••	Saturated solution	BaCls
5	••	• •	• •	• •	• •
6	• •	• •	• •	none	
7	• 3	••	••	Saturated solution	BaCls
8	박yodak micro-mesh	••	••	••	• •
11	• •	• •	••	Saturated solution	CaCla
12	American Colloid Volclay	••	••	••	• 4
13	Wyodak micr o-mesh	••	••	none	
14	American Colloid Volclay	••	••	Saturated solution	Ba(OH) _s
17	Wyodak micro-mesh	• •	• •	Saturated solution	Ca (OH)2
19	••	••	••	Saturated curous nit solution	

•	Subsequent	Treatme	en t	reensoidT (eadou)	Dielectric Gonstant	Power Factor	Number
•	Heated at for 2 hour			0.005	6.98	11.1(per	cent) 1
	Hested st			0.005	6.42	3.3	2
	none	A1 3		0.003	21.35	81.77	3
	none			0.003	10.6	16.4	4
	Rested at			0.0023	9.4	12.9	5
	• •	••	• •	0.0015	4.82	5.6	6
1	Heated at 17000 lbs			0.002	9.1	10:6	7
'	Hested at	560 C;	2 hr.	0.002	7.40	4.07	8
	Heated at	560 C;	14 hrs.	0.005	7.43	11.9	11
	••	••	••	0.003	6.19	27.87	12
	Heated at	560 C;	14 hrs.	0.005	33,15	63 • 8	13
	••	••	• •	0.005	4.76	5 .78	14
	••	••	••	0.0035	5.25	2.03	17
•	••	••	••	0.003	7.47	1.3	19

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4	Subsectiont Tre	etment	Thickness (inches)	Dialoctric Constant	Power Factor	dumber	
	Hested at 560 14 hrs.	for	0.003	9.64	57.9 (percent) 20	
	Heated at 184 17000 lbs. pro	C; l hr.,	0.00175	7.01	2.3	21	
	•• , ••	••	0.0015	8.44	11.9	22	
,	••	••	0.00175	9.92	68.6	23	
· .	Heated at 560 14 hrs.	C for	0.002	6.41	12.5	101	
	••	••	0.0035	11.96	46.8	102	
	••	••	0.010	20.6	••••	103	
	••	••	0.0015	8.38	83.5	104	
	••	••	0.0045	5.34	7.58	106	
	none		0.015	5.30	17.1	112	

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TABLE VII (Continued)

Sample Number	Clay Used	Remarks on Film Propagation	Base Exchange Agent
20	Wyodak micro-mesh	Clay suspension poured to 2 mm. thickness, then dried at 50 C in cabinet	Saturated mercuric nitrate solution
51	••	••	Saturated lead acetate solucion
22	••	••	Satura ted mercurous nitrate solution
23	••	••	Saturated mercuric nitrate solution
101	2 •	Film formed from colloidal [el of clay containing a 1.2 percent soln. of BaClg	none
102	••	Film formed from colloidal (el of electrodialyzed clay	••
103	••	Same as 101 except that a 10 percent soln. of BaCl _a was used.	••
104	American Colloid Volclay	This film formed by eir drying at R. T. for 48 hrs.	••
105	••	Same as 104 but dried in ovan at 50 C.	••
112	Fisher Sentonite		• •

Subsequent Trestment Heated at 560 C for 14 hours	Thickness (inches) 0.0025	Dielootric Constant 3.93	Fower Fector 34.36	Number 201
••	0.004	9.85	29.6	301
••	0 .0 035	7.64	2.8	302
••	0.006	5.31	5.4	303
••	0.004	4.99	9.15	304
••	0.003	3.12	5.4	307
••	0.0015	1.29	81.4	308
••	0.002	1.59	57.4	310

TABLE VII (Concluded)

Semple Number	Clay Used	Clay suspension Oured to 2 mm. thickness, then dried at 50 C. in a cabinet		Bese Exchange Agent Seturated carium nitrate solution
201	Wyodak micro-mesh			
301	••	••	••	Saturated sodium stannate solution
302	••	• •	••	Saturated CoClassolution
303	••	• •	••	Saturated cadmium scetate solution
304	••	• •	••	Saturated Cr ₂ (30 ₄), solution
307	••		••	Saturated Th(NO ₃) ₄ solution
308	••	• •	••	Seturated aluminum potassium sulfate solution
310	• •	• •	••	Saturated uranium acetate solution

of one hour and then a continued heating at this temperature for a period of 14 additional hours. The film strips, cut three inches by two inches, were blaced between class plates to keep them from curling. A Lindberg furnace was employed. It appeared to be critical that the initial rise in temperature to 550 degrees Centigrade be not too rapid. With rapid increases in temperature the film would be reduced to dust as it was dehydrated. When the treatment was carried out in the regular way the films became black in color and were much more brittle than they were before the curing step.

TESTING OF THE FILM FOR DIBLECTRIC CONSTANT AND POWER FACTOR

A GENERAL RADIO Type 716-C Caparithmee Bridge with Type 1302-A Oscillator and Type 1231-B Amplifier and Mull Detector was used in the testing procedure. A mercury electrode holder specially made according to a model submitted by Mr. F. H. Wells (16) was employed to hold the film strip. The electrode had a one square inch surface area contact between both sides of the 111m and the mercury which connected in turn with shielded leads to the bridge itself.

Before the test of a film, the film was kept in a drying oven at 100 degreer Centigrade for at least 24 hours. The test was made never more than 15 minutes after the strip was removed from the oven. The electrode holder was kept in a desicoator until the time of the test. Under such conditions, daily humidity fluctuations did not seem to be of much importance. A film which was tested two hours after removal from the oven would have increased noticeably in power factor.

The properties and average test results of the various films

prepared are liven in Table VII.

RESULTS

An analysis of the data in Table VII indicates several interesting facts. Wyodak films seem to be superior to American Colloid films if the base exchange and subsequent treatment has been identical. Compare semples 1 and 8, 11 and 12 for values of the dielectric constant and the power factor. It is of interest to see the differences in chemical composition of these two clays. See Table II.

Samples 11 and 17 seemed to indicate that the use of the free base of the metal is to be preferred to the use of the saturated salt of the metal although the salt is definitely much more soluble in each case.

From samples 1 and 2 it appeared that longer periods of heating in the furnace greatly improved the power factor with only slight reduction in the dielectric constant.

In a few cases increase in molecular weight of the cation in the base exchange reaction seemed to cause the power factor of the film to decrease (samples 8, 11 and 19). In many other cases that relation did not hold true. A consistent but unexplained difference existed between the effect of mercurous and mercuric cations in the base exchange (samples 19, 20, 22, and 23).

Substitution of pressures up to 17000 lbs. per sq. in., shorter periods of heating and lower temperatures in place of the 560 degree Centigrade and 24 hour curing period did not give comparable results, as far as the power factor was concerned (contrast samples 19 and 22, 2 and 7). However, a one hour period at 175 degrees Centigrade and 17000 lbs. per sq. in. seemed to be more favorable than a ten minute treatment under the same conditions (see samples 5 and 7).

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